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APPLICATION NO.	FII	LING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO	
09/758,902	09/758,902 01/11/2001		Roberts S. David	PC9047D	1327	
23913	7590	590 11/28/2005		EXAM	EXAMINER	
PFIZER INC	C		DUFFY, PATRICIA ANN			
150 EAST 42 5TH FLOOR			ART UNIT	PAPER NUMBER		
NEW YORK			1645			

DATE MAILED: 11/28/2005

Please find below and/or attached an Office communication concerning this application or proceeding.

			ication No.	Applicant(s)	Applicant(s)				
	Office Astinu Commence	09/7	58,902	DAVID ET AL.	DAVID ET AL.				
Office Action Summary			niner	Art Unit					
			cia A. Duffy	1645					
Period fo	The MAILING DATE of this commun or Reply	ication appears o	n the cover sheet	with the correspondence a	ddress				
WHIC - Exte after - If NC - Failu Any	ORTENED STATUTORY PERIOD F CHEVER IS LONGER, FROM THE M nsions of time may be available under the provisions SIX (6) MONTHS from the mailing date of this common period for reply is specified above, the maximum state to reply within the set or extended period for reply reply received by the Office later than three months are ded patent term adjustment. See 37 CFR 1.704(b).	IAILING DATE O of 37 CFR 1.136(a). In nunication. atutory period will apply will, by statute, cause the	F THIS COMMU no event, however, may and will expire SIX (6) May be application to become	NICATION. y a reply be timely filed NONTHS from the mailing date of this ABANDONED (35 U.S.C. § 133).					
Status									
1)	Responsive to communication(s) file	ed on <i>27 Januar</i> v	2005.						
2a)□	•	2b)⊠ This action							
3)	Since this application is in condition for allowance except for formal matters, prosecution as to the merits is								
·	closed in accordance with the practice under <i>Ex parte Quayle</i> , 1935 C.D. 11, 453 O.G. 213.								
Disposit	ion of Claims				•				
4)⊠)⊠ Claim(s) <u>18-20</u> is/are pending in the application.								
	4a) Of the above claim(s) is/are withdrawn from consideration.								
5)	Claim(s) is/are allowed.								
6)⊠	Claim(s) 18-20 is/are rejected.								
7)	Claim(s) is/are objected to.								
8)[Claim(s) are subject to restrict	ction and/or electi	ion requirement.						
Applicat	ion Papers								
9)[The specification is objected to by th	e Examiner.							
10)[The drawing(s) filed on is/are	a) accepted	or b) Objected	to by the Examiner.					
	Applicant may not request that any obje	ction to the drawing	g(s) be held in abe	yance. See 37 CFR 1.85(a).					
	Replacement drawing sheet(s) including			T .					
11)	The oath or declaration is objected to	by the Examine	r. Note the attacl	ned Office Action or form P	PTO-152.				
Priority (ınder 35 U.S.C. § 119								
	Acknowledgment is made of a claim ☐ All b) ☐ Some * c) ☐ None of:	for foreign priority	y under 35 U.S.C	C. § 119(a)-(d) or (f).					
	Certified copies of the priority documents have been received. Certified copies of the priority documents have been received in Application No								
	3. Copies of the certified copies	of the priority do	cuments have be	en received in this Nationa	al Stage				
	application from the Internation	nal Bureau (PCT	Rule 17.2(a)).						
* (See the attached detailed Office action	on for a list of the	certified copies r	ot received.					
Attachmen	· ·								
_	ce of References Cited (PTO-892) ce of Draftsperson's Patent Drawing Review (F	PTO-048\		w Summary (PTO-413) No(s)/Mail Date					
3) Infor	mation Disclosure Statement(s) (PTO-1449 or No(s)/Mail Date		~ ·	of Informal Patent Application (PI	ΓΟ-152)				

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DETAILED ACTION

The response after final filed 1-27-05 has been entered into the record. Claims 1-17 have been canceled and claims 18-20 are pending and under examination.

The finality of the rejection of the last Office action is withdrawn in view of the new grounds of rejection set forth below.

Applicants should note that the examiner in charge of this application has changed.

Please address all future correspondence to Exr. Patricia A. Duffy.

Sequence Requirements

This application contains sequence disclosures that are encompassed by the definitions for nucleotide and/or amino acid sequences set forth in 37 C.F.R. § 1.821(a)(1) and (a)(2). However, this application fails to comply with the requirements of 37 C.F.R. § § 1.821-1.825 for the reason(s) set forth below. Full compliance with the sequence rules is required in response to this office action.

Double Patenting

The nonstatutory double patenting rejection is based on a judicially created doctrine grounded in public policy (a policy reflected in the statute) so as to prevent the unjustified or improper timewise extension of the "right to exclude" granted by a patent and to prevent possible harassment by multiple assignees. See *In re Goodman*, 11 F.3d 1046, 29 USPQ2d 2010 (Fed. Cir. 1993); *In re Longi*, 759 F.2d 887, 225 USPQ 645 (Fed. Cir. 1985); *In re Van Ornum*, 686 F.2d 937, 214 USPQ 761 (CCPA 1982); *In re Vogel*, 422

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F.2d 438, 164 USPQ 619 (CCPA 1970); and, In re Thorington, 418 F.2d 528, 163 USPQ 644 (CCPA 1969).

A timely filed terminal disclaimer in compliance with 37 CFR 1.321(c) may be used to overcome an actual or provisional rejection based on a nonstatutory double patenting ground provided the conflicting application or patent is shown to be commonly owned with this application. See 37 CFR 1.130(b).

Effective January 1, 1994, a registered attorney or agent of record may sign a terminal disclaimer. A terminal disclaimer signed by the assignee must fully comply with 37 CFR 3.73(b).

Claims 18-20 are rejected under the judicially created doctrine of obviousness-type double patenting as being unpatentable over claims 1-21 of U.S. Patent No. 6,083,512 in view of Farmers and Consumers Market Bulletin (Department of Agriculture, Atlanta, Georgia, 70(24): 1984, page 1, 12, ill.) and .

The claims of the patent teach all of the elements of the claimed invention except for the inclusion of a viral immunogen.

Farmers and Consumers Market Bulletin disclose a clostridial vaccine composition which comprises a viral immunogen from influenza, equine viral rhinopneumonitis, strangles and teaches the annual vaccination with the multivalent vaccine reduces the threat of infection from both the bacterium and the virus.

Kensil et al show the use of a saponin adjuvant in association with an antigen, wherein the exemplified vaccine comprises a viral antigen. Kensil establishes the adjuvant activity of saponin is effective for viruses.

It would have been *prima facie* obvious to one having ordinary skill in the art at the time that the invention was made to include any of the respiratory viral immunogens of Farmers and Consumers Market Bulletin in the multicomponent Clostridial vaccines of the patent because Kensil et al teach that saponins are effective for adjuvanting a viral

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antigen and Farmers and Consumers Market Bulletin disclose the conventional combination of Clostridial vaccine antigens with respiratory viral antigens.

Claim Rejections - 35 USC § 103

The following is a quotation of the appropriate paragraphs of 35 U.S.C. 102 that form the basis for the rejections under this section made in this Office action:

The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains.

Patentability shall not be negatived by the manner in which the invention was made.

This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(e), (f) or (g) prior art under 35 U.S.C. 103(a).

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Claims 18 and 20 are rejected under 35 U.S.C. 103(a) as being unpatentable over Seifert (Deutsche Tiearzliche Wochem. 90(7):274-279, 1983) in view of Geresi et al (Ann. Immunol. Hung. 25(0):37-40, 1985), Farmers and Consumers Market Bulletin (Department of Agriculture, Atlanta, Georgia, 70(24): 1984, page 1, 12, ill.), and Kensil (US Patent 5,057,540 issued 1991).

Seifert teaches the use of a saponin adjuvant in the formulation of a multivalent clostridial vaccine using toxins of apparently different strains of Clostridial pathogens for the purposes of obtaining enhanced protective immune responses in a host. Seifert teach that the group has isolated three local pathogens from anaerobic infections and these pathogens are used for producing the anatoxin. The anatoxin from the three pathogens are toxoided with formalized sodium chloride solution and the toxoids are purified and concentrated. The toxoid vaccine is mixed with anthrax spores in saponin and administered intracutaneously at a dose of 0.2 ml/animal, twice at an interval of 4 weeks. The vaccine provided for a marked protective effect (see page 2). Seifert et al differ by not providing for the addition of antigens from a respiratory virus and multiple serotypes/species.

Geresi et al teach the formulation of multivalent clostridial vaccine compositions Clostridium perfringens antigens (C and D-type toxins: different serotypes) and tetanus toxoid (different species), which also comprise a viral antigen (see page 38).

Farmers and Consumers Market Bulletin disclose a clostridial vaccine composition which comprises a viral immunogen from influenza, equine viral rhinopneumonitis, strangles and teaches the annual vaccination with the multivalent vaccine reduces the threat of infection from both the bacterium and the virus.

Kensil et al show the use of a saponin adjuvant in association with an antigen, wherein the exemplified vaccine comprises a viral antigen. Kensil establishes the adjuvant activity of saponin is effective for viruses.

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It would have been *prima facie* obvious to one having ordinary skill in the art at the time that the invention was made to modify the Seifert vaccine by adding any desired additional clostridial components as taught by Geresi or Farmers and Consumers Market Bulletin and include a respiratory viral antigen as taught by Farmers and Consumers Market Bulletin because Geresi and Farmers and Consumers Market Bulletin teach that it is conventional to combine the multivalent clostridial vaccine with viral components and both Seifert and Kensil teach the use of saponin as an effective adjuvant for the enhancement of an immune response with either a clostridial or viral antigen respectively the combined vaccine would provide the advantage of reduced time and cost for administering multiple vaccines to farm/ranch animals. Absent unexpected results, one of skill in the art would expect the modified composition to protect from infection because the saponin adjuvant was effective to generate protective immunity in cattle and that similar compositions with different adjuvants were also effective to generate protective immunity.

Claim 19 is rejected under 35 U.S.C. 103(a) as being unpatentable over Seifert (Deutsche Tiearzliche Wochem. 90(7):274-279, 1983), Geresi et al (Ann. Immunol. Hung. 25(0):37-40, 1985), Farmers and Consumers Market Bulletin (Department of Agriculture, Atlanta, Georgia, 70(24): 1984, page 1, 12, ill.), and Kensil (US Patent 5,057,540 issued 1991) as applied to claims 18 and 20 above further in view of Green et al (The Veterinary Record, 120:435-439, 1987).

The teachings of Seifert (Deutsche Tiearzliche Wochem. 90(7):274-279, 1983) in view of Geresi et al (Ann. Immunol. Hung. 25(0):37-40, 1985), Farmers and Consumers Market Bulletin (Department of Agriculture, Atlanta, Georgia, 70(24): 1984, page 1, 12, ill.), and Kensil (US Patent 5,057,540 issued 1991) as combined are set forth above. The teachings differ by not explicitly combining immunogens from six or more species or serotypes of *Clostridium*.

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Green et al teach the formulation of a multi-valent clostridial vaccine for the purposes of stimulating a protective immune response against multiple serotypes and species of this pathogen. Green et al teach three known commercially available vaccines comprising at least 7 different serotypes/species of Clostridium for protection from infection (Tasvax, Heptavac, Covexin) page 435, column 2, Table 1.

It would have been *prima facie* obvious to one having ordinary skill in the art at the time that the invention was made to modify the composition as combined *supra* by adding the other known individual clostridial vaccine toxoid components of Green et al (i.e. C. perfringens (serotype D), C. septicum, C. novi, C. haemolyticum and C. chauveoi) because combined vaccines were commercially available and known to be effective for broad protection for a variety of pathogens in farm animals. Given the demonstrated efficacy and commercial availability of the 7- and 8- way combination vaccines, the combination as combined would be expected to be effective to protect from infection.

Status of the Claims

All claims stand rejected.

Conclusion

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free).

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Any inquiry concerning this communication or earlier communications from the examiner should be directed to Patricia A. Duffy whose telephone number is 571-272-0855. The examiner can normally be reached on M-Th 6:30 am - 6:00 pm. If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Lynette Smith can be reached on 571-272-0864.

The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Patricia A. Duffy, Ph.D.

Primary Examiner

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